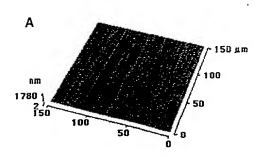
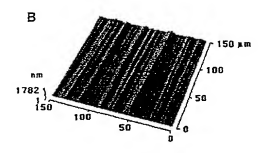
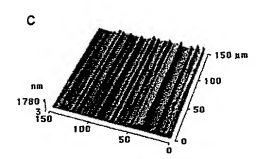


Oriented NFG Substrates: Comparison of Surface Topography







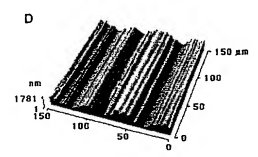


Figure 2 3

Atomic Force Microscopy Images of Oriented NFG Substrates

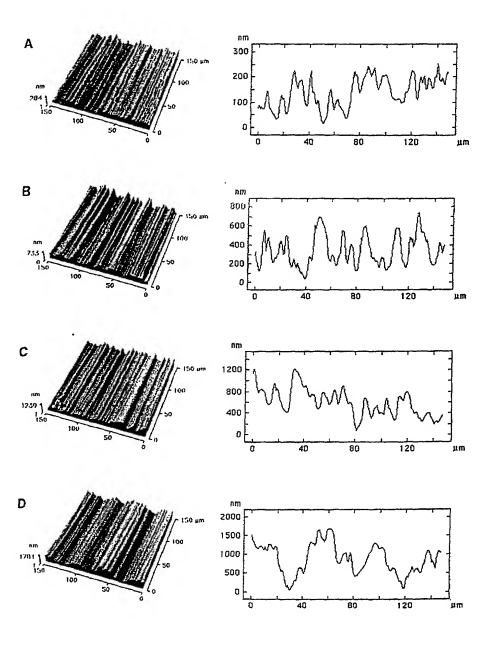
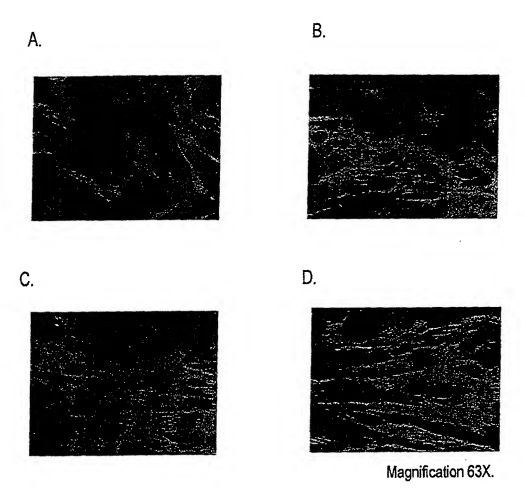


Figure 3 4

Meningeal Cytoskeletal Alignment on Oriented NFG Substrates



FITC-labeled microtubules (red) and DAPI stained nuclei. Cell orientation is lost on surfaces with a surface roughness of less than 1000nm.

Figure 45

Primary Meningeal Cells on Polypropylene of Various Oriented Microtopography

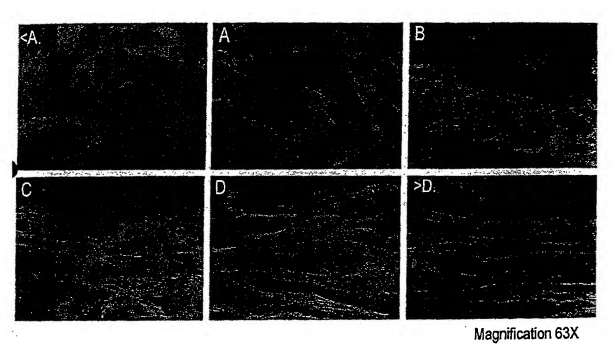
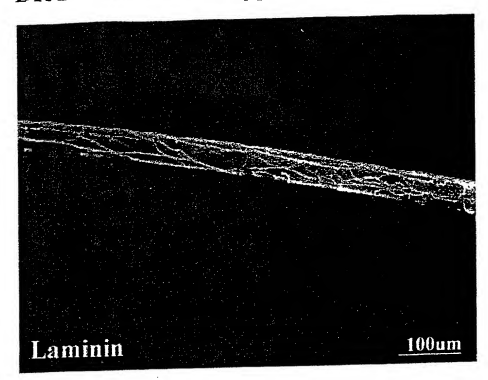
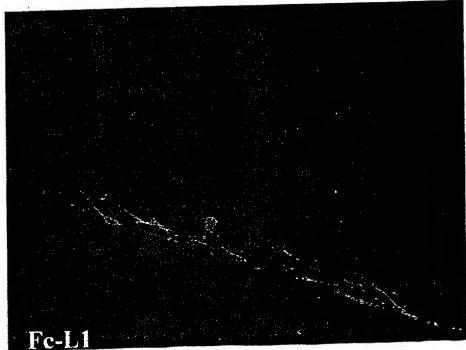


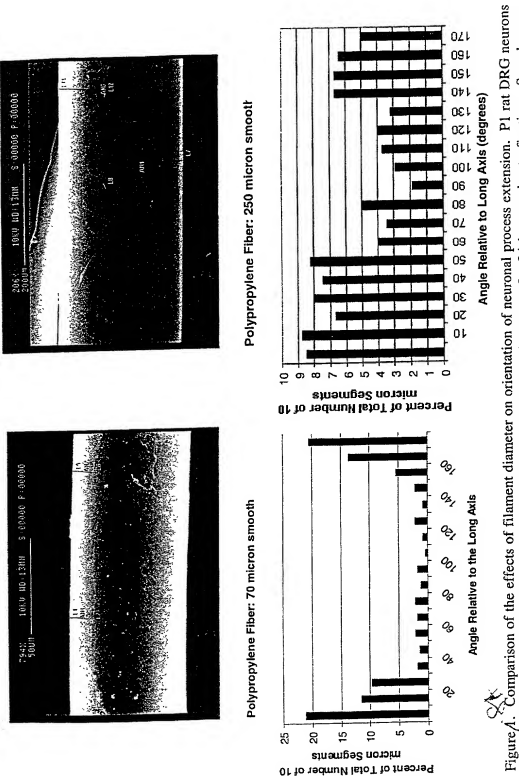
Figure \$ C

DRG Neurons on Polypropylene Filaments



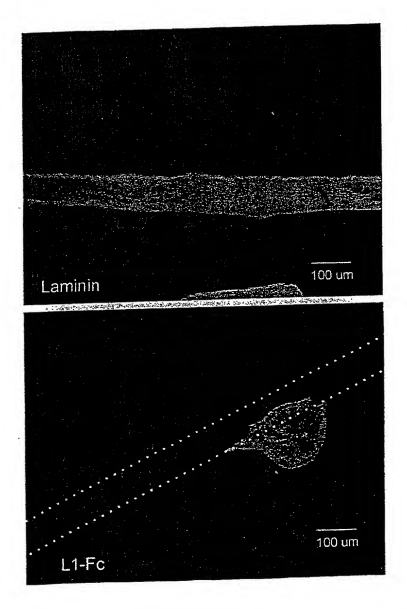


FIGURT.

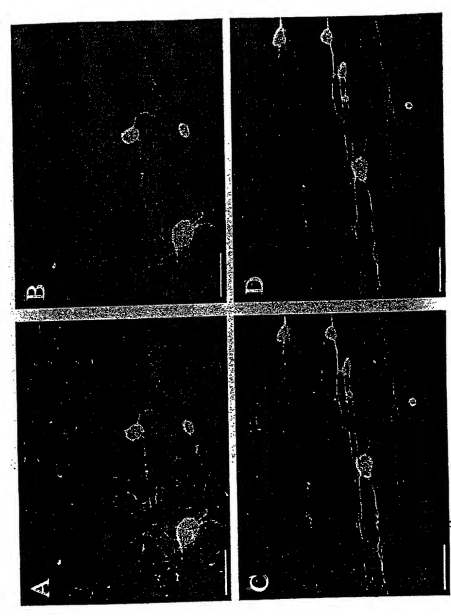


were plated on laminin-treated polypropylene filaments and allowed to grow for 24 hours prior to fixation. Sample were processed by standard methods for scanning electron microscopy. Representative images and a summary of data are shown for two filament sizes.

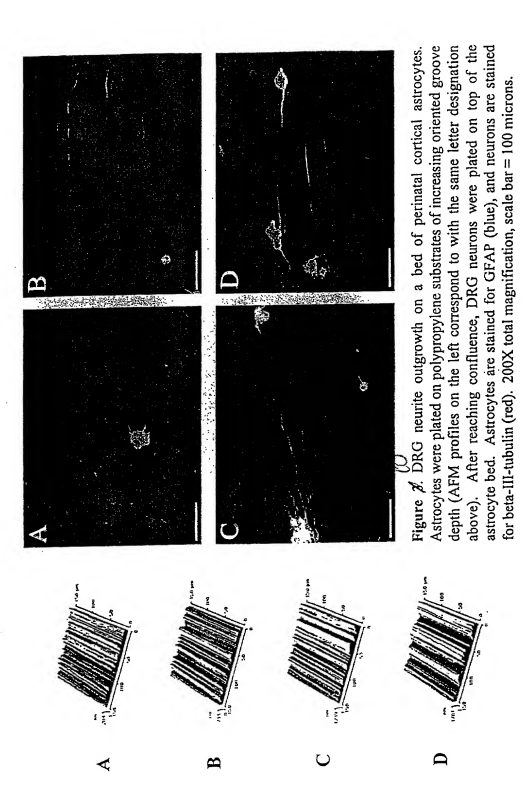
Astrocyte Attachment to Polypropylene: Effect of Laminin and L1-Fc

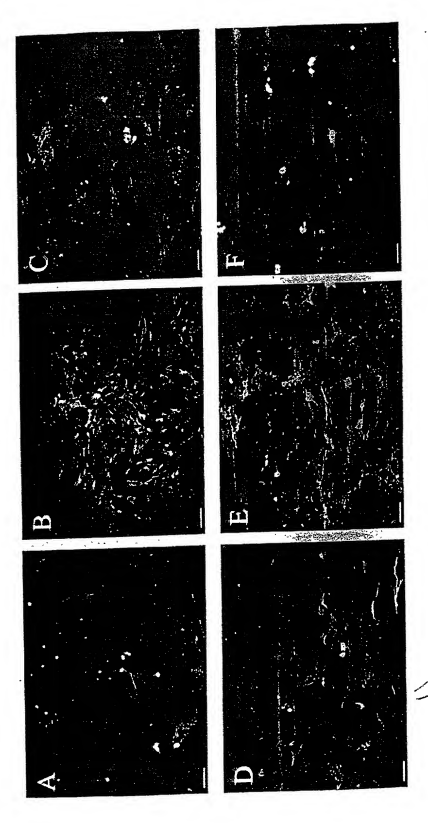




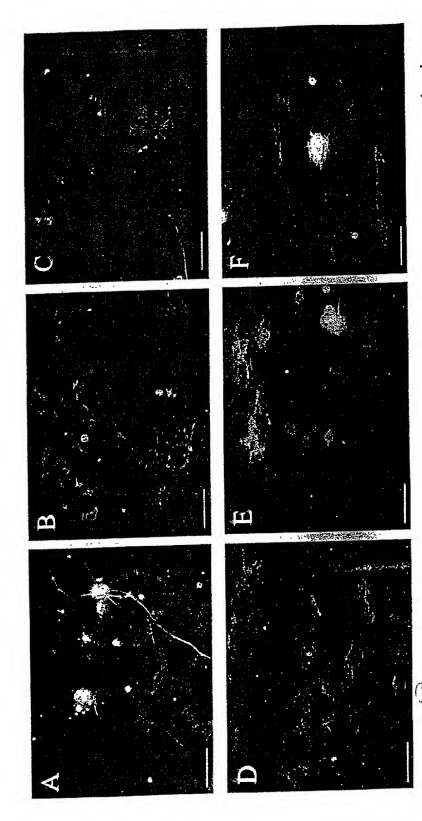


and panel D depicts the same field as C. In both B and D, the green channel for fibronectin has been omitted to enhance the contrast of the beta-III-tubulin stain. Total magnification is substrates. Panels A and B depict the smoothest surface, while panels C and D show a highly grooved surface. Meningeal cells are labeled with cellular fibronectin (green), neurons with beta-III-tubulin (red), and all nuclei with DAPI (blue). Panel B depicts the same field as A, Figure X'A comparison of neurite outgrowth on meningeal cells grown on grooved 200X, scale bar = 100 microns.





through F (A=smoothest and F=roughest). 100X total magnification, scale bar = 100 microns. A higher polypropylene substrates of increasing groove depth. The underlying astrocytes were stained for actin Astrocytes were plated on (red). Neurons were stained for beta-III-tubulin (green). Oriented groove depth increases from panel A Figure A. DRG outgrowth on a bed of perinatal cortical astrocytes. power of view of the same conditions is shown on the next figure.



polypropylene substrates of increasing groove depth. The underlying astrocytes were stained for actin (red). Neurons were stained for beta-III-tubulin (green). Oriented groove depth increases from panel A through F (refer to corresponding AFM images of substrates). 200X total magnification, scale bar = 100 Figure 8. DRG outgrowth on a bed of perinatal cortical astrocytes. Astrocytes were plated on microns.

FIGURES:

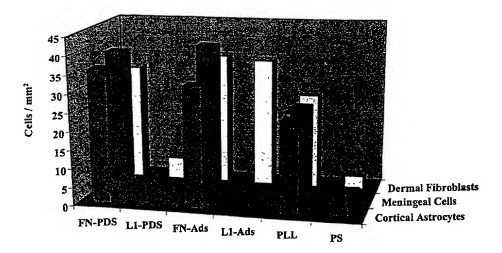


Figure X. Comparison of cell attachment to substrates treated with fibronection, L1, and poly-1-lysine. The following surface treatments were compared: fibronectin covalently bound to polyethylene oxide (PEO) (FN-PDS); L1 covalently bound to polyethylene oxide (PEO) (L1-PDS); fibronectin adsorbed to the substrate (FN-ADs); L1 adsorbed to the substrate (L1-ADs); poly-1-lysine adsorbed to the substrate (PLL); and nothing adsorbed to the substrate (PS). All cultures were maintained in the presence of serum containing media.

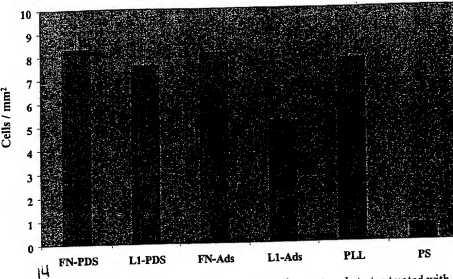


Figure 2 Comparison of dorsal root ganglion cell attachment to substrates treated with fibronection, L1, and poly-1-lysine. The following surface treatments were compared: fibronectin covalently bound to polyethylene oxide (PEO) (FN-PDS); L1 covalently bound to polyethylene oxide (PEO) (L1-PDS); fibronectin adsorbed to the substrate (FN-ADs); L1 adsorbed to the substrate (L1-ADs); poly-1-lysine adsorbed to the substrate (PLL); and nothing adsorbed to the substrate (PS). All cultures were maintained in the presence of serum containing media. Attachment to L1-Ads and PS were significantly

less than other groups.

groups.

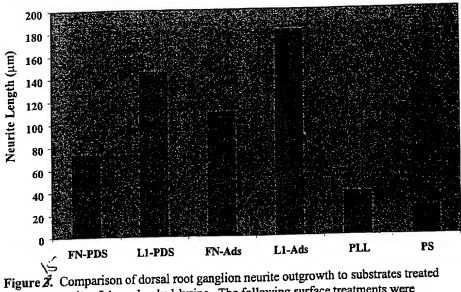


Figure 3. Comparison of dorsal root ganglion neurite outgrowth to substrates treated with fibronection, L1, and poly-l-lysine. The following surface treatments were compared: fibronectin covalently bound to polyethylene oxide (PEO) (FN-PDS); L1 covalently bound to polyethylene oxide (PEO) (L1-PDS); fibronectin adsorbed to the substrate (FN-ADs); L1 adsorbed to the substrate (L1-ADs); poly-l-lysine adsorbed to the substrate (PLL); and nothing adsorbed to the substrate (PS). All cultures were maintained in the presence of serum containing media. Outgrowth on both L1 treatments were significantly greater that FN treatments which were also greater that PLL and PS

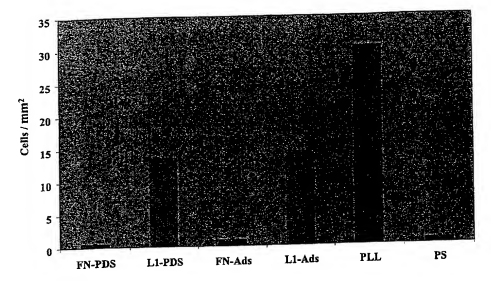


Figure . Comparison of cerebellar granule neuron cell attachment to substrates treated with fibronection, L1, and poly-1-lysine. The following surface treatments were compared: fibronectin covalently bound to polyethylene oxide (PEO) (FN-PDS); L1 covalently bound to polyethylene oxide (PEO) (L1-PDS); fibronectin adsorbed to the substrate (FN-ADs); L1 adsorbed to the substrate (L1-ADs); poly-1-lysine adsorbed to the substrate (PLL); and nothing adsorbed to the substrate (PS). All cultures were maintained in the presence of serum containing media. Attachment to L1-treatments and PLL were significantly greater than other groups.

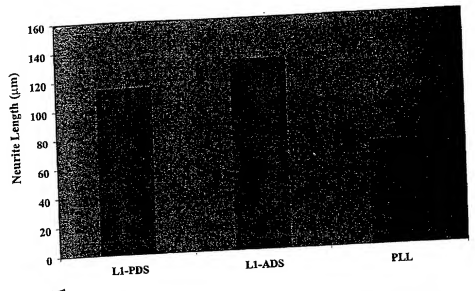


Figure 5. Comparison of cerebellar granule neuron neurite outgrowth on substrates treated with L1 and poly-1-lysine. The following surface treatments were compared: fibronectin covalently bound to polyethylene oxide (PEO) (FN-PDS); L1 covalently bound to polyethylene oxide (PEO) (L1-PDS); fibronectin adsorbed to the substrate (FN-ADs); L1 adsorbed to the substrate (L1-ADs); poly-1-lysine adsorbed to the substrate (PLL); and nothing adsorbed to the substrate (PS). All cultures were maintained in the presence of serum containing media. No outgrowth occurred on the FN or the PS conditions (data not shown). Outgrowth on both L1 treatments were significantly greater PLL.

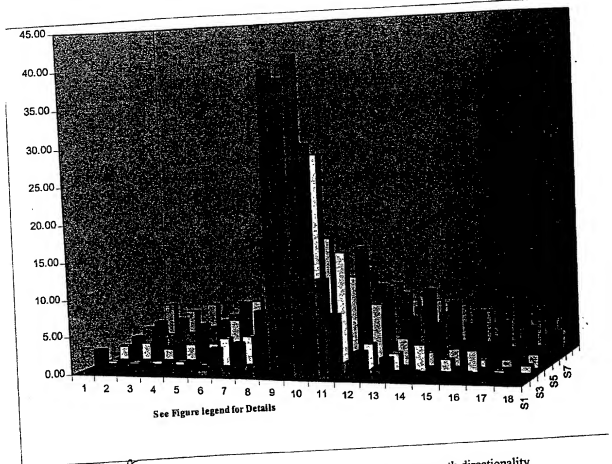
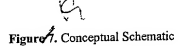
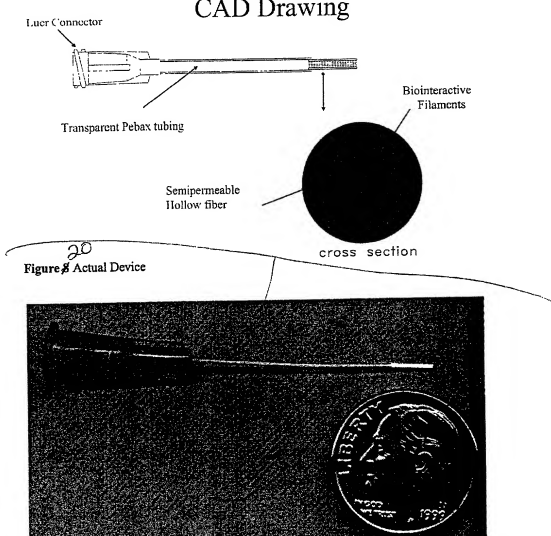


Figure 4. Comparison of primary dorsal root ganglion neurite outgrowth directionality on polypropylene substrates with differing radius of curvature. The following surfaces were diameters (microns): S1 was 53; S2 was 95; S3 was 200; S4 was 300; S5 was 500; and S6 was a flat surface. Data is reported as the percentage of outgrowth in angles segment of 10 degrees where 1=0; 2=10; 3=20;...10=90 or growth perpendicular to the segment of no on. On-going studies will fill in the data by examining smaller diameter long axis; and so on. On-going studies will fill in the data by examining smaller diameter filaments and by increasing the number of observations at each filament diameter so that a model can be created that predicts cell behavior as a function of filament a diameter.



Multi-filament NFG device: CAD Drawing



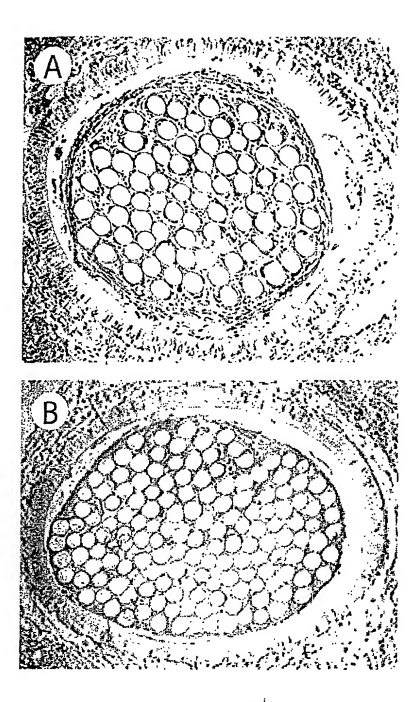


Figure 77 21

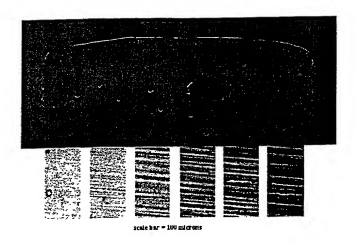


Figure \(\beta \) Low power light micrograph of polymeric substrate, with varying degrees of microtexture, used for cell culture studies.

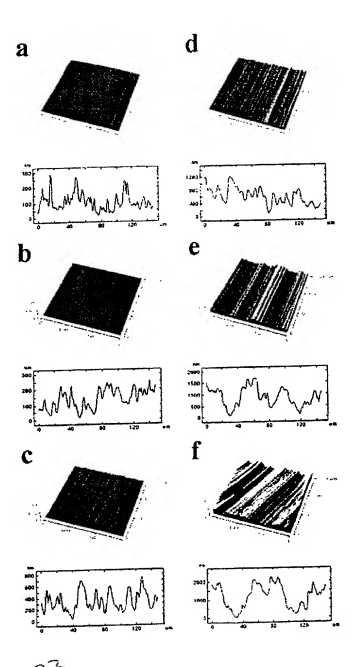


Figure 9 Representative images of varying surface topography of substrate shown in figure 8 as determined by scanning atomic force microscopy.

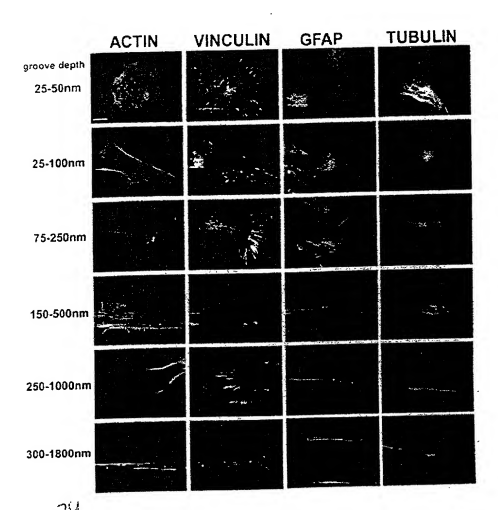


Figure 10 Gradual alignment of the astrocyte cytoskeleton as a function of increasing groove depth in the surface topography. 100x immunofluorescence photomicrographs of cells stained for a range of cytoskeletal proteins. Scale bar = 20 microns.

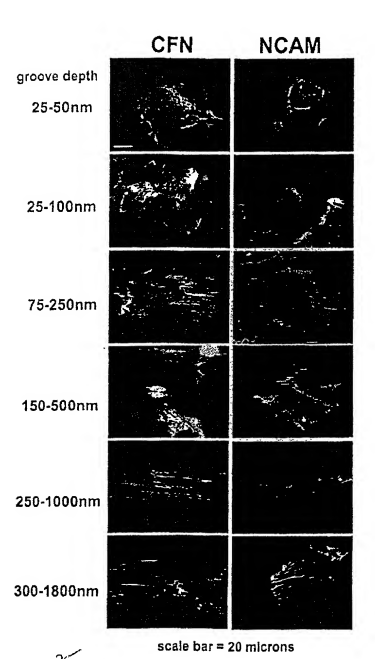


Figure 17 Gradual alignment of astrocyte-derived extracellular matrix and cell adhesion proteins as a function of increasing groove depth. 100x immunofluoresescence photomicrographs of cells stained for CFN or NCAM.

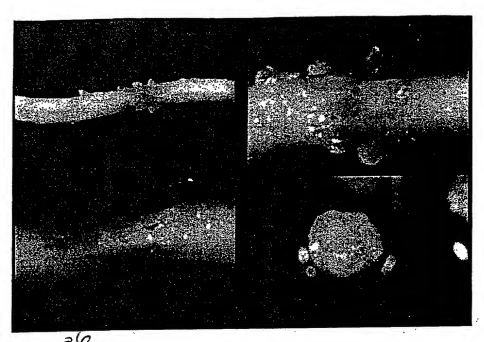
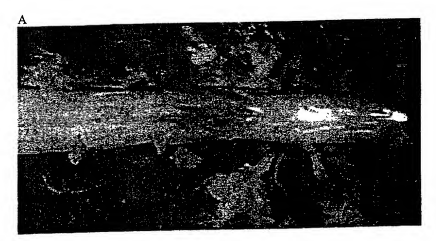


Figure 12 Photographs of rat thoracic spinal cord with dorsal hemisection and implantation of a multi-filament device. The integration of the device is shown from dorsal view before (above) and after (bottom left) removal of the dura. The presence of the device within the dorsal lesion space can be seen in transverse section in the bottom right.



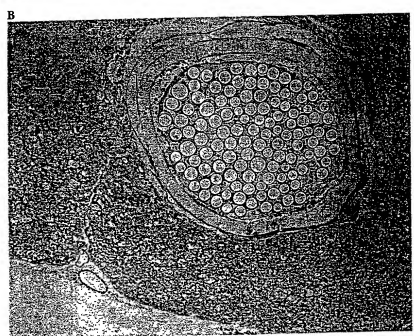


Figure 13 A) Dorsal view of the spinal cord following fixation for histology and exposure within the vertebral canal. The site of injury and implantation of the multi-filament device is clearly visible, though the reaction to the implant is minimal.

B) Transverse section through the multifilament device within the spinal cord, just outside the injury zone. The filaments within the semi-permeable tube are clearly visible, surrounded by cellular material that has completely invaded the lumenal space of the device. Luxol fast blue/H & E stain.

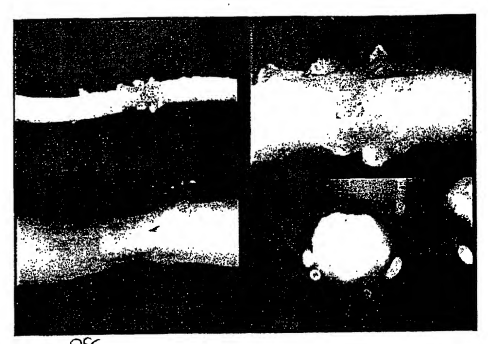


Figure 12 Photographs of rat thoracic spinal cord with dorsal hemisection and implantation of a multi-filament device. The integration of the device is shown from dorsal view before (above) and after (bottom left) removal of the dura. The presence of the device within the dorsal lesion space can be seen in transverse section in the bottom right

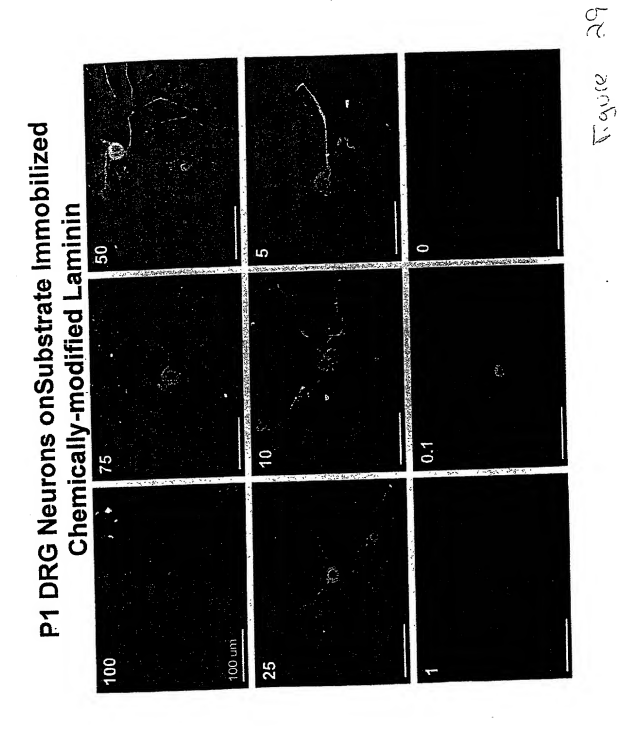


Figure 30

P1 DRG Neurons on Substrate Immobilized Chemically-modified Fibronectin

